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- 6) A marked up copy of the specification;
- 7) Letter to Draftsperson with Formal Figures; and,
- 8) A receipt indication postcard.

AMENDMENT

Please amend the specification and claims as follows.

In the Specification.

Delete the paragraph at page 15, lines 13-15 and insert the following:

Figures 6A-6F illustrate the use of the *blm* NRPS and PKS enzymes to synthesize a variety of hybrid polyketide/peptide molecules including, but not limited to, a family of oxazolines/oxazoles, and thiazoline/thiazoles. Figure 6A synthesis using BlmIX, BlmVIII, and BlmVII. Figure 6B synthesis using NRPS, BlmVIII, and BlmVII. Figure 6C synthesis using BlmIX, BlmVIII, and NRPS (C, A^N, PCP). Figure 6E synthesis using BlmIX, BlmVIII and NRPS (C, A^C, PCP). Figure 6F synthesis using BlmIX, BlmVIII, and NRPS (C, A^C, PCP, OX).

Delete the paragraph at page 19, lines 12-19 and insert the following:

The nucleic acids comprising the *blm* gene cluster are identified in Tables I and II and listed in the sequence listing provided herein (SEQ ID NOS: 1 and 2, GenBank Accession number AF210249 (which replaces sequence AF149091), and SEQ ID NO:3, GenBank Accession number AF210311). In particular, Table I identifies genes and functions of open reading frames (ORFs) responsible for the biosynthesis of the hybrid peptide/polyketide/peptide backbone and sugar moieties of bleomycin, while Table II identifies a number of ORFs comprising the *blm* gene cluster, identifies the activity of the catalytic domain encoded by the ORF and provides primers for the amplification and isolation of that orf.

Delete the Table 1 at pages 19-20 and insert the following:

Table I. Determined functions of ORFs in the bleomycin biosynthesis gene cluster

Gene	Amino acids	Sequence Homolog'	Proposed function ^{2, 3}
orf8	424 SEQ ID NO: 115	YqeR (BAA12461)	Oxidase







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orf9 (blmC)	498	RfaE (AAD07904)	NDP-glucose synthas€ENTER 160
	SEQ ID NO: 114		
orf10 (blmI)	90	GrsB (P14688)	Type II PCP
	SEQ ID NO: 113		
orf11 (blmD)	545	NodU (Q53515	Carbamoyl transferase
,	SEQ ID NO: 112		
orf12 (blmE)	390	RfaF (AAD16056)	Glycosyl transferase
), j = (0)	SEQ ID NO: 111		
orf13	187	MbtH (O05821)	Unknown
71,113	SEQ ID NO: 110		
orf14 (blmII)	462	Nrp (CAA98937)	NRPS condensation enzyme
71j1 4 (0titi1)	SEQ ID NO: 109	Imp (et al bess)	1.12 2 30.120.131.101.101.101.101
ouf15	339	SyrP (1890776)	Regulation
orf15	SEQ ID NO: 108	3911 (1050770)	Regulation
ouf16 (blIII)	<u> </u>	HMWP2 (P48633), McbC	A PCP Ox
orf16 (blmIII)	935	(P23185)	
orf17 (hlmIV)	SEQ ID NO: 107 2626	HMWP2 (P48633)	C A PCP Cy A PCP Cy
orf17 (blmIV)	SEQ ID NO: 106	1111111 2 (1 40033)	
orf18	638	AsnB (2293165)	Asparagine synthetase
<i>31</i> J10	SEQ ID NO: 105	11311D (2273103)	7 isparagine symmetase
orf19 (blmF)	494	RfbC (Q50864)/BlmOrf1	Glycosyl transferase/β-
orgio (bunii)	SEQ ID NO: 104	(507319)	hydroxylase
orf20 (blmG)	325	YtcB (2293288)	Sugar epimerase
onjeo (ounis)	SEQ ID NO: 103	1102 (22/5200)	July of the state
orf21 (blmV)	645	McyB (2708278)	PCP C
3. J2 1 (3)	SEQ ID NO: 102	120,2 (2,002,0)	
orf22 (blmVI)	2675	ACoAS (1658531), PksD	A ⁴ ACP C A PCP C A
3 (- ···· · - /	SEQ ID NO: 101	(S73014)	
	020111111111111111111111111111111111111	SnbDE (CAA67249)	
orf23 (blmVII)	1218	SyrE (3510629)	C A PCP
, ,	SEQ ID NO: 100		
orf24 (blmVIII)	1841	HMWP1 (CAA73127)	KS AT MT KR ACP
•	SEQ ID NO: 99		
orf25 (blmIX)	1066	SafB (1171128)	C A PCP
	SEQ ID NO: 98		
orf26 (blmX)	2162	TycC (2623773)	C A PCP C A PCP
	SEQ ID NO: 97		
orf27 (blmXI)	688	SyrE (3510629)	NRPS condensation enzyme
	SEQ ID NO: 96		
orf28	239	SC9C7.04C (CAA22716)	Unknown
<u>-</u>	SEQ ID NO: 95		
orf29	582	YvdB (CAB08068)	Transmembrane transporter
	SEQ ID NO: 94		
orf30	113	SmtB (P30340)	Regulation
<u> </u>	SEQ ID NO: 93		
orf31	117	PhnA (P16680)	Unknown
	SEQ ID NO: 116		



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Delete Table II at pages 21-23 and insert the following replacement Table II:

Orf #	Activity	Method	and primers for ORF amplifi	Seq
011#	Activity	Memod	Forward	ID
			Reverse	No.
orf-8	Oxygen-independent	Gapped-blast	F: ATGAGCCACGCCATCGGA	5
SEQ ID NO:115	coproporphyrinogen	comparison	R: TCAGGCGCGTTCGGGGGC	6
3EQ ID NO.113	III oxidase	Comparison	n. Tenedecerreces	*
orf-9	ADP-heptose synthase	Gapped-blast	F: GTGAACACCGACCTGCCC	7
SEQ ID NO:114	(blmC)	comparison ¹	R: TCATGGGGTGTCTCCCTC	8
orf-10	Peptidyl carrier	Expression and	F: ATGAGCGCCCCGCGGGC	9
SEQ ID NO:113	protein	biochemical	R: TCACCGGTCCCGCTCCCC	10
	(blmI)	characterization. ²		
orf-11	Carbamyltransferase	Gapped-blast	F: ATGAGCGCCGACCCGTCC	11
SEQ ID NO:112	(blmD)	comparison ¹	R: TCATGAGCGGGCCGCCGT	12
orf-12	ADP-heptose:LPS	Gapped-blast	F: ATGACCACCCCATGACC	13
SEQ ID NO:111	heptosyl transferase (blmE)	comparison ¹	R: TCATGGGGTACTCCTGAT	14
orf-13	Homolog of mbtH in	Gapped-blast	F: ATGACCACGACCCCGCGG	15
SEQ ID NO:110	the synthesis of	comparison ¹	R: TCAGGTGCCGGACACGCG	16
-	mycobactin			ļ
orf-14	Peptide synthetase	Gapped-blast	F: GTGACCGCCCCGGCACA	17
SEQ ID NO:109	(condensation, blmII)	comparison ¹	R: TCATCGGTGGCTCCTCGT	18
orf-15	Regulatory gene	Gapped-blast	F: GTGAACCGGCACGGCCCC	19
SEQ ID NO:108	(homolog of syrP)	comparison ¹	R: TCACGCGCTCACCTCGTC	20
orf-16	Mutated peptide	Gapped-blast	F: GTGACGAGCGCCCGGCCC	21
SEQ ID NO:107	synthetase- oxidase (NRPS-0, blmIII)	comparison ¹	R: TCACGGGGCCTCCGTGCG	22
orf-17	Peptide synthetase	Expression and	F: ATGCTGCACGGCGCCGCG	23
SEQ ID NO:106	(NRPS-2-1,blmIV)	biochemical	R: TCACTCCGGTCCACCTCC	24
520121101	(2.22.0.2.2,0)	characterization. ²		
orf-18	Asparagine synthetase	Gapped-blast	F: GTGAGGCCCGTGTGCGGC	25
SEQ ID NO:105		comparison ¹	R: TCAGCCACCGTTGCCGCC	26
orf-19	Homolog of	Gapped-blast	F: GTGAAGGACCTCGGCCGG	27
SEQ ID NO:104	hydroxylase-	comparison ¹	R: TCACTCCCCGGTGCCGG	28
	dehydrogenase (blmF)			
orf-20	Nucleotide-sugar	Gapped-blast	F: GTGACATGGACCGTGGTG	29
SEQ ID NO:103	epimerase (blmG)	comparison ¹	R: TCAGGCATCGGCCCTCCC	30
orf-21	Peptide synthetase	Gapped-blast	F: ATGCGCGGGCATGACGAC	31
SEQ ID NO:102	(NRPS-3CT, blmV)	comparison ¹	R: TCACGGTGTCTCTCCCTC	32
orf-22	Peptide synthetase	Expression and	F: ATGAGCCGGCCGGCC	33
SEQ ID NO:101	(NRPS-5-4-3, blmVI)	biochemical	R: TCATGCTCGGTCATCGCC	34
		characterization. ²		
orf-23	Peptide synthetase	Expression and	F: GTGACCACGCCCGCATC	35
SEQ ID NO:100	(NRPS-6, blmVII)	biochemical characterization. ²	R: TCATTCGGGACGCGGCA	36
orf-24	Polyketide synthase	Gapped-blast	F: ATGAGCCATGCCGACGCG	37
SEQ ID NO:99	(blmVIII)	comparison ¹	R: TCACAGCACCACCTCTTC	38
orf-25	Peptide synthetase	Gapped-blast	F: ATGACCCCGGCCGCCGAC	39
SEQ ID NO:98	(NRPS-7, blmIX)	comparison	R: TCATCGTCCGCCGCCTTT	40
orf-26	Peptide synthetase	Gapped-blast	F: ATGCCTCGGTGTGCCCGA	41
SEQ ID NO:97	(NRPS-9-8, blmX)	comparison ¹	R: TCATTCGGCGGCACCTCC	42



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orf-27	Peptide synthetase	Gapped-blast	F: GTGGGTTTCCGTCGAGCG R: TTACACCCTCCGTTTCTC	43 44
SEQ ID NO:96	(condensation, blmXI)	comparison ¹		45
orf-28	Phosphatidylserine	Gapped-blast	F: ATGGCACAGGACCTGAAC	45
SEQ ID NO:95	decarboxylase	comparison ¹	R: TCAACGCCACCGGATCTT	
orf-29	Transmembrane	Gapped-blast	F: GTGAGCTCCCTCGCCGTC	47
SEQ ID NO:94	transporter	comparison ¹	R: TCATCGTCGGGCACTCGG	48
orf-30	Metal dependent	Gapped-blast	F: GTGCCGGTTCCGCTGTAT	49
SEQ ID NO:93	regulatory element	comparison ¹	R: TCACCGGGCACTGACCTC	50
orf-31	PHNA homolog	Gapped-blast	F: GTGACCGAGAACCTTCCG	51
SEQ ID NO:116		comparison	R: TCAGACCTTCTTGACCAC	52
orf-32	Peptide synthetase	Gapped-blast	F: ATGGCCTCAGACGCTTTG	53
SEQ ID NO:117	(NRPS-11-10)	comparison ¹	R: TCATTGAGACTCCTCCTC	54
orf-33	Putative transporter	Gapped-blast	F: ATGATGAAGTCAAGCCGC	55
SEQ ID NO:118		comparison	R: TCAGTGGCTTACAAGGAG	56
orf-34	Homolog of	Gapped-blast	F: ATGACTGACCTGCCGTTG	57
SEQ ID NO:119	clavaminic acid	comparison ¹	R: TCACACCAGCAGCGAGGT	58
	synthase			
orf-35	Thioesterase	Gapped-blast	F: ATGGATTTCCCCCTCACC	59
SEQ ID NO:120		comparison ¹	R: TCATGCCCCTACCTCGGC	60
orf-36	Putative transporter	Gapped-blast	F: ATGACCGCGCGCGTCGAC	61
SEQ ID NO:121		comparison ¹	R: TCACTCCTCGGCTTCGGC	62
orf-37	Unknown	Gapped-blast	F: GTGTCCAAGAACGCGGCG	63
SEQ ID NO:122		comparison ¹	R: TCATCGGCTCGCCTCGTG	64
orf-38	Peptide synthetase	Gapped-blast	F: ATGACCCTCACCCTGCGG	65
SEQ ID NO:123	(NRPS-12)	comparison ¹	R: TCACTCGGGCACTCCTTC	66
orf-39	Regulatory gene	Gapped-blast	F: GTGACCGGTTCCGTAACG	67
SEQ ID NO:124	(homolog of SyrP	comparison ¹	R: TCATGAGTCCGCCGAGGT	68
orf-40	Peptide synthetase	Gapped-blast	F: ATGACAGAGGTCCGAGGT	69
SEQ ID NO:125	' '	comparison ¹	R: CCCGGCAACCGCCCTCCC	70
orf-41	4'-	Expression and	F: GTGATCGCCGCCCTCCTG	71
SEQ ID NO:126	phosphopantetheinyl	biochemical	R: TTACGGGACGGCGGTCCG	72
	transferase (pptA)	characterization. ²]

Delete the paragraph on page 69, line 17 through page 70, line 20 and insert the following:

The sequence of the 1,761-bp BamHI-SalI fragment was analyzed for coding regions by using the CODONPREFERENCE and TESTCODE programs of the GCG package (Genetics Computer Group, Madison, Wisconsin). Two complete ORFs (pptA, orf3) and two incomplete ORFs (orf1, orf4) were identified within the sequenced region (Figure 13). The first ORF from left to right (designated orf1) starts out of the analyzed area and ends with a TGA codon at position 248 of the sequenced fragment. Comparison of the deduced product of orf1 with proteins encoded by nucleic acids in databases showed similarities with Rv2795c from Mycobacterium tuberculosis (GenBank AL008967) and SC5A7. 22 from S. coelicolor (GenBank AL031107), both of unknown function. The second ORF, pptA, contains the sequence amplified by PCR and used for the cloning of this locus. It comprises 741 nucleotides, starting with a GTG codon (position 245) which is





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coupled to the stop codon of orf1, and ending with a TAA codon. The starting codon of pptA is preceded by a potential ribosomal binding site (RBS), GGGAG. The overall (76.6%) and third codon position (93. 9%) G+C contents and the codon usage of pptA are similar to those found in other Streptomyces genes, with the exception of the stop codon (TAA), which is most uncommon in this group of organisms (Wright et al. Gene (1992) 113:55-65). The pptA gene encodes a protein of 246 amino acids with a predicted molecular mass of 25,619 Da and a pI of 4. 76, which contains the conserved PPTase motifs. Databases searches with PptA showed significant similarities to the putative actinomycete PPTases (39-52%/48-61% identity/similarity) and to confirmed bacterial PPTases such as EntD from E. coli (17%/24% identity/similarity) (Lambalot et al. Chem. Biol. (1996) 3:923-936). The third ORF, orf3, is separated from pptA by an apparently noncoding DNA region of 153 bp, and it is transcribed in opposite and convergent direction with respect to orf1-pptA. The gene orf3 comprises 240 nucleotides, starting with an ATG codon (position 1358) and ending with TGA. The starting codon of orf3 is preceded by the sequence GAAGG, a potential RBS. The deduced product of orf3 encodes a protein of 79 amino acids with a predicted mass of 7,555 Da and a pI of 7. 17. The Orf3 protein shows similarities to the N-terminal region of SC5H1. 35c, a protein of unknown function from S. coelicolor (encoded by nucleic acid sequence in GenBank AL049863). Analysis of Orf3 with the SignalP program (Nielsen et al. *Protein Engineer*. (1997) 10:1-6) predicts an N-terminal signal peptide which would be cleaved between residues 27 and 28 (ALA-DS), suggesting that the mature protein (52 amino acids, 5,099 Da, pI 4. 31) would be secreted. Between orf3 and orf4 there is an apparently noncoding region of 251 nucleotides. The orf4 gene is transcribed in opposite and divergent direction with respect to orf3. It starts with an ATG codon at position 1610, preceded by a potential RBS (GGAGG), and ends out of the sequenced fragment. The deduced protein product (50 amino acids) of the incomplete orf4 contains a potential NAD/FAD binding motif, GXGX₂GX₃GX₆G (SEQ ID NO:92) (Scrutton et al. *Nature* (1990) 343:38-43), showing low similarities to diverse oxidoreductases.

In accordance with 37 CFR §1.121 a marked up version of the above-amended paragraph(s) illustrating the changes introduced by the forgoing amendment(s) are provided in Appendix C.

